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CORROSION OF CHEMICALLY VAPOUR DEPOSITED TITANIUM CARBIDE ON AN INERT SUBSTRATE

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Abstract - The corrosion properties of chemically vapour deposited titanium carbide (TiC) have been studied. Any contribution to the corrosion current from the substrate was avoided by using an inert substrate.

The TiC was deposited onto substrates of Al₂O₃ from a gas mixture containing TiCl₄, CH₄ and H₂ at temperatures between 1020°C and 1080°C. Differences in film thickness and morphology were measured for samples deposited at different temperatures and gas flows. Film thicknesses varied between 0.5 and 1.5 μm and grain sizes between 0.1 and 0.8 μm.

Anodic polarization curves were registered in 1 M hydrochloric acid (HCl). HCl of a lower concentration did not activate the samples. The corrosion potentials of the samples were used as starting points for the polarization curves. Films deposited on polished Al₂O₃ at 1020°C reached stable potentials of 250 mV (vs. SCE) after one hour. Films deposited onto unpolished substrates had lower values of corrosion potentials (~100 to 0 mV vs. SCE) which reached stability after about three hours. It is suggested that the passivating film on TiC, deposited on polished substrates, achieved a higher protective quality in the solution, due to self-healing. The passivating film on TiC, deposited on unpolished substrates, on the other hand, did not succeed to reach this high quality.

Anodic polarization curves for the different films showed only slight differences. The curves showed two passivation peaks at 800 and 1900 mV (vs. SCE), respectively. An extremely high oxygen overpotential (>5V) and a poor tendency to transpassive corrosion (<6V vs. SCE) were indicated by the curves.

INTRODUCTION

Titanium carbide thin films are used for many technical applications. Cutting and punching tools and high wear bearings coated with TiC have been used because of its high hardness. Films of TiC have also found use as redox electrodes on various substrates and as adhesion improving interlayers. TiC deposition for corrosion protection has been used in the chemical processing industry and in waste water treatment plants [1].

Increasing effort has been made to optimize the properties of deposited (CVD or PVD) films by varying deposition parameters [2,3]. Much work has been done to determine the mechanical properties of TiC films [4,5] and some investigations have also been made on their corrosion properties [6]. There is always a possibility, however, that the substrate contributes to the corrosion behaviour and thus reliable corrosion data on the film itself can not be measured. However, such data can be obtained by depositing the film on an inert substrate. This has previously been done for TiC deposited onto quartz substrates by activated reactive evaporation [7]. In this work the inherent corrosion properties of CVD TiC is reported.
EXPERIMENTAL

Films of TiC were deposited onto both unpolished and polished polycrystalline substrates of Al₂O₃ (>99.5%) (see table 1). Polishing was done with diamond paste (15 µm). Prior to the deposition all the samples were cleaned in an ultrasonic bath with freon. The Al₂O₃ substrates were then loaded into a hot wall CVD reactor and, during the heating of the furnace, hydrogen (99.9997%) was allowed to flow over them. Deposition took place at 1020 to 1080°C and a total pressure of 50 torr (see table 1). When reaction temperature was reached, the gas mixture containing 9% methane (99.9995%), 6% TiCl₄ (99.999%) and 85% H₂ (99.9997%) was allowed to flow through the reactor for one to two hours.

When the desired reaction time had passed the gas flows and furnace were shut off and only hydrogen was allowed to flow through the system during cooling. Upon reaching room temperature, the hydrogen flow was turned off and argon (99.9997%) was admitted into the reactor while removing the samples.

Table 1. CVD parameters for the three batches.

<table>
<thead>
<tr>
<th>Batch name</th>
<th>Temp. °C</th>
<th>Gas flow velocity sccm</th>
<th>Dep. time h</th>
<th>Polished substrates</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>1080</td>
<td>200</td>
<td>1</td>
<td>None</td>
</tr>
<tr>
<td>B</td>
<td>1020</td>
<td>200</td>
<td>2</td>
<td>None</td>
</tr>
<tr>
<td>C</td>
<td>1020</td>
<td>100</td>
<td>2</td>
<td>All</td>
</tr>
</tbody>
</table>

Phase content and texture coefficients were determined with X-ray diffractometry (Philips PW 1710). From experimental values of lattice parameter vs. carbon content /8/, the carbon content in the films could be estimated. The morphology of the samples were studied in a scanning electron microscope (SEM, JEOL-840). One sample from each experiment was then cut off and the film thickness measured in the SEM. Film thicknesses were also calculated from quantitative analysis with X-ray fluorescence spectroscopy (XRFS).

The samples were glued onto copper cylinders with silver glue to create an electric contact between the sample and the rotating electrode and cold mounted in an epoxy resin (Struers epofix), so that only the sample surface could be exposed to the electrolyte. Before the corrosion testing took place the samples were cleaned with acetone in an ultrasonic bath.

The electrolytes were prepared from concentrated HCl of analytical grade and deionized water. The resulting solution was deaerated by one hour of water aspiration followed by nitrogen purge during the corrosion measurements. The corrosion cell was thermostatted to 24°C. The samples could not be activated in 0.1 M HCl. However, in 1 M HCl the samples were activated. With a PAR 273 potentiostat the corrosion potential of each sample was recorded for three hours. It had then reached stability. The anodic polarization curve was now recorded up to 6V (vs. the saturated calomel electrode (SCE)), with a scanning rate of 0.1 mV per second and the speed of the rotating electrode being 3000 rpm.

After the corrosion measurements, the epoxy and the copper cylinders were removed from the samples. The samples were analysed again with X-ray diffraction, SEM and XRFS.

RESULTS

The X-ray diffractograms from the uncorroded samples showed peaks from TiC (Khaembraevit, JCPDS-file 32-1383) and Al₂O₃ (Corundum, JCPDS-file 10-173). Texture coefficients (TC) were calculated by using equation 1 for the different reflecting angles of TiC.
\[ TC_{hkl} = \frac{I'_h k l / I_{h k l}}{\frac{1}{n} \sum_{i=1}^{n} (I'_{h_i k_i l_i} / I_{h_i k_i l_i})} \]

\( I' \) is the measured intensity, \( I \) is the calculated intensity, \( hkl \) are the indices for the reflections and \( n \) is the number of reflecting angles.

The [200] direction dominated the A and B batches (\( TC_{200} = 1.3 \)), whereas the [113] direction dominated the C batch (\( TC_{113} = 1.6 \)).

Table 2 shows measured values of grain sizes and film thicknesses for the samples. Grain sizes varied between 0.1 and 0.8 \( \mu \text{m} \), the lowest values being for the C batch and the highest for the A batch. Film thicknesses of uncorroded samples vary between 0.5 and 1.5 \( \mu \text{m} \).

Table 2. Grain sizes and film thicknesses of the batches

<table>
<thead>
<tr>
<th>Batch</th>
<th>Grain size/( \mu \text{m} )</th>
<th>Film thickness/( \mu \text{m} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>0.5-0.8</td>
<td>1-2</td>
</tr>
<tr>
<td>B</td>
<td>0.2-0.6</td>
<td>1-2</td>
</tr>
<tr>
<td>C</td>
<td>0.1-0.5</td>
<td>0.5-1</td>
</tr>
</tbody>
</table>

The open circuit potential was measured for three hours and stable potential values in the range -100 to 350 mV (vs. SCE) were reached. The lower values represent the A and B batches which in some cases reached an unstable higher potential value after about 1.5 hours and which then rapidly dropped. Potential time curves are shown in fig.1.

Fig.1 Measured potential vs. time (at zero current).
Surfaces of uncorroded samples were grey and TiC on polished substrates had a metallic lustre. The surfaces of corroded samples were dark grey. For the C samples, the corroded film was very thin and had a light grey colour. SEM of corroded samples displayed attacked areas in grain boundaries and that sharp, triangular grains in A and B are rounded.

The anodic polarization curves of the three different batches are shown in fig. 2. Two active peaks at 800 and 1800 mV (vs. SCE), respectively appear. Critical current density \( i_{\alpha} \) for the first active peak ranges from 30 to 570 \( \mu \text{A-cm}^{-2} \) and passive current density \( i_p \) between 2 and 47 \( \mu \text{A-cm}^{-2} \). For the second peak \( i_{\alpha} \) is 1 to 7 mA-cm\(^{-2}\) and \( i_p \) varies between 25 and 190 \( \mu \text{A-cm}^{-2} \). After the second passive region the current density does not increase again but decreases or remains constant at least up to 6V (vs. SCE), where the recording was ended, and no transpassive corrosion occurred. No oxygen evolution could be detected, indicating that the oxygen overpotential was higher than 5 volts.

X-ray diffractograms for corroded samples showed only peaks of TiC and Al\(_2\)O\(_3\). No peaks from any corrosion products could be found.

**DISCUSSION**

Variations in deposition parameters related to batch characteristics and corrosion behaviour.
Samples prepared in the C batch yielded the thinnest films. For the C batch, a lower gas flow velocity than in the B batch was employed. Furthermore, polished substrates were used in the C batch and not in the B batch. Theoretically one would expect a higher deposition rate with a lower gas flow. However, Lindström and Amberg have shown that, for CVD of TiC and at sufficient supply of reactants, the deposition rate is not influenced by the total gas flow rate. The only remaining explanation is that the deposition rate decreases with the polishing of a substrate. On an unpolished substrate there will be a higher density of nucleation sites, which might explain the thicker films obtained in batch B.

No clear distinction of corrosion behaviour between the samples can be seen in the polarization curves. Measurement of corrosion potential vs. time, however, shows a decided difference between the C batch and the two other batches. As can be seen in Fig. 1, potential values of C samples reach 200 to 250 mV (vs. SCE) after about one hour and remain stable for the rest of the recording time. Potential values of A and B samples, on the other hand, either remain low with slight variations or reach a higher value only to make a sudden drop to the lower values where they remain.

It has been suggested in earlier work that spontaneous passivation of TiC occurs even at very low oxygen concentrations. The reason for the high open circuit potential could therefore be that the cathodic reaction is a sum of O₂-reduction and hydrogen evolution. The easiest explanation for the higher value of the steady state potential for the C batch is higher oxygen concentration in these solutions. The solutions have, however, been treated identically in all experiments and there is still an apparent difference in corrosion potential between the C batch on one hand and the A and B batches on the other.

A rise in corrosion potential is the result of either increased anodic polarization or decreased cathodic polarization. (Decreased cathodic polarization would be the case in a more oxygen-rich solution.) When a passivating film is formed on the surface it forms a barrier through which reactants diffuse with difficulty. Evans measured the potential of a metal with a discontinuous oxide film /11/. His experiments showed that breakdown of the film resulted in potential drops whereas a self-healing film increased the potential. In another work /12/ Bannister and Evans conclude that a high final potential, at zero current, indicates immunity from attack. It should be noted that the corrosion potential discussed here, gives no information about the corrosion current or corrosion rate. Two plausible explanations as to why the samples show differences in corrosion potentials will be given here.

It has been documented that specific solutions etch different crystal planes at different rates /13/. The preferred orientations vary between the batches (see "Results") and this might give differences in the rate at which they passivate.

Another, perhaps more significant, explanation for these discrepancies is the difference in surface smoothness due to polishing of the substrates. A deposit with a smooth surface will, upon oxidation, form a stronger and continuous protective layer. The higher values of steady state potential for the C samples, as well as the shorter time of getting there, could thus be explained by the formation of better protecting films.

2. Comparison with Pourbaix diagram.

It may be fruitful to discuss the anodic polarization curves in relation to a Pourbaix diagram. The Pourbaix diagram for TiC can be studied in Fig. 3. The dotted line in the diagram indicates pH 0. According to mixed potential theory the corrosion potential should lie between the reversible potentials of H⁺/H₂ and TiC/Ti⁵⁺ (-244 and -420 mV vs. SCE, respectively). The recorded values of 10 to 350 mV (vs. SCE), being higher than the reversible potential of H⁺/H₂, indicates residual oxygen in the electrolyte and spontaneous passivation. According to the diagram the stable phase at the first active peak is TiO₂·H₂O and the reaction here should be one of the following:

\[
\text{TiC} + 4\text{H}_2\text{O} \leftrightarrow \text{TiO}_2\cdot\text{H}_2\text{O} + \text{CO(g)} + 6\text{H}^+ + 6\text{e}^-(2)
\]

\[
\text{TiC} + 5\text{H}_2\text{O} \leftrightarrow \text{TiO}_2\cdot\text{H}_2\text{O} + \text{CO}_2\text{(g)} + 8\text{H}^+ + 8\text{e}^-(3)
\]

When the potential is increased and the current once again rises, the oxidation of TiO₂·H₂O to TiO₃·2H₂O takes place according to equation (4):

\[
\text{TiO}_2\cdot\text{H}_2\text{O} + 2\text{H}_2\text{O} \leftrightarrow \text{TiO}_3\cdot2\text{H}_2\text{O} + 2\text{H}^+ + 2\text{e}^-(4)
\]
3. Related works

Hintermann and Cowling have made a thorough investigation of the corrosion products of CVD TiC in sulphuric acid [15]. The TiC was deposited onto steel discs which were then removed and polarization experiments were conducted on the remaining flakes of TiC (which contained up to four percent each of iron and chromium). The recorded polarization curve of TiC in 2M H₂SO₄ shows an active peak at about 1050 mV (vs. SHE) corresponding to 830 mV (vs. SCE), which is in good agreement with this work. At 1800 mV (vs. SHE) further dissolution begins but no peak value can be determined since the recording ends at 2000 mV (vs. SHE).

Corrosion products at the first peak were determined in galvanostatic, coulometric and rotating ring disc electrode experiments. The authors concluded that the dissolution of TiC produced TiO₂·H₂O, with Ti³⁺ as an unstable intermediate, and CO, which was either desorbed or oxidized to CO₂. Above 970 mV (vs. SHE) CO is said to be the stable carbon species. Presuming that corrosion products are the same in H₂SO₄ and HCl, equation 2, above, should be the valid one for the present experiment. For the second current rise it was assumed that TiO₂²⁺ is formed which can not be true for our case, except as an intermediate reaction, since passivity is once again reached.

Morancho et al. [10] registered anodic polarization curves of polycrystalline sintered TiC. They reported a corrosion potential in 5N sulphuric acid solutions of 420 mV (vs. SHE) corresponding to 180 mV (vs. SCE) which is almost the mean value for the present work. Polarization curves showed one peak at 1150 mV (vs. SHE) and a broad one at 2200 to 2600 mV (vs. SHE). These values are higher than those in the present work which might be attributed to the fact that a lower pH was used by Morancho (pH=-0.7).
Alloca et al. /16/ reported corrosion potentials of MOCVD-TiC in 0.1 N H₂SO₄ of 300 to 400 mV (vs. SCE). They reported only a slight difference between values in aerated solutions and solutions deaerated by bubbling purified nitrogen into the system. Since titanium and its divalent ion easily oxidises and there is always a small amount of oxygen left in the solution after deaeration with either water aspiration or nitrogen purge the “true” value of the corrosion potential of TiC in deaerated solutions might be very hard to measure. The first active peak of our experiments at 800 mV (vs. SCE) is not detected by the authors which might be a result of the high scanning rate, 111 mV/min, compared with 6 mV/min in the present work.

Beverskog et al. /7/ performed corrosion measurements on TiC which had been deposited onto quartz by activated reactive evaporation (ARE). The polarization curves were recorded with the same experimental set-up as in the present work. There is only one active peak in the diagram and at a lower value than for the CVD TiC (710 mV vs. SCE). At approximately 1800 mV (vs. SCE) there is a slight current increase which might indicate the oxidation of TiO₂.H₂O to TiO₃.2H₂O as shown in eq.4. The ARE TiC evidently passivates easier, as shown by the lower values of passivation potential and also by the fact that the second passivation peak is only implied. Values of critical current density and passivation current density (60 μA.cm⁻² and 25 μA.cm⁻² respectively) agree well with values in the present work.

CONCLUSIONS

The inherent corrosion properties of hot wall CVD TiC, deposited at 1020 to 1080°C on polished and unpolished Al₂O₃ have been determined. Corrosion potentials in 1M HCl varied between -100 and 350mV (vs. SCE). When recording the polarization curve two current density peaks, at 800 and 1800 mV (vs. SCE), respectively, could be registered. It is presumed that the first peak corresponds to the oxidation of Ti²⁺ to Ti³⁺ which then immediately reacts to form TiO₂.H₂O (see /13/). The second peak is due to the oxidation of TiO₂.H₂O to TiO₃.2H₂O. Critical current density (i₉) for the first active peak ranges from 30 to 570 μA.cm⁻² and passive current density (i₉) lies between 2 and 47 μA.cm⁻². For the second peak i₉ is 1 to 7 mA.cm⁻² and i₉ varies between 25 and 190 μA.cm⁻². The TiC showed no tendency to transpassive corrosion (<6V vs. SCE) and had a high oxygen overpotential (>5V).

Potential-time curves of the uncorroded samples show discrepancies between the batches. TiC deposited onto polished Al₂O₃ (C samples) show rising potentials which stabilize, at about 250 mV (vs. SCE), after about one hour. TiC deposited onto unpolished Al₂O₃ (A and B samples), on the other hand, have unstable corrosion potentials up to three hours, where they reach stable values of -100 to 0 mV (vs. SCE). This discrepancy between the samples is explained by a more protective oxide film on the C samples.

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